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(54) Title: DELTA 6 FATTY ACID DESATURASE

(57) Abstract

Novel human DNA sequences that encode the gene CYB5RP, a delta 6 fatty acid desaturase, are provided. Provided are genomic CYB5RP DNA as well as cDNA that encodes the CYB5RP protein. Also provided is CYB5RP protein encoded by the novel DNA sequences. Methods of expressing CYB5RP protein in recombinant systems are provided. Also provided are CYB5RP methods that identify activators and inhibitors of CYB5RP protein.

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TITLE OF THE INVENTION DELTA 6 FATTY ACID DESATURASE

CROSS-REFERENCE TO RELATED APPLICATIONS Not applicable.

STATEMENT REGARDING FEDERALLY-SPONSORED R&D Not applicable.

10 REFERENCE TO MICROFICHE APPENDIX Not applicable.

FIELD OF THE INVENTION

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The present invention is directed to novel human DNA sequences encoding a delta 6 fatty acid desaturase, an enzyme involved in the synthesis of essential fatty acids.

BACKGROUND OF THE INVENTION

Essential fatty acids (EFAs) are polyunsaturated fatty acids that cannot be manufactured by mammals, yet are required for a number of important biochemical processes, and thus must be supplied in the diet. The most important dietary EFAs are linoleic acid and alpha-linolenic acid (ALA). These two EFAs undergo a number of biosynthetic reactions that convert them into various other EFAs. Figure 1 depicts the biosynthetic reactions involving the two groups of EFAs, the n-6 EFAs (linoleic acid derivatives) and the n-3 EFAs (ALA derivatives). EFAs are formed from linoleic acid and ALA by a series of alternating reactions involving the removal of two hydrogens coupled with the insertion of an additional double bond (desaturation) and the lengthening of the fatty acid chain by the addition of two carbons (chain elongation). The enzymes catalyzing the desaturations and elongations are thought to be the same for both groups of EFAs.

Among the more important unsaturated fatty acids are the delta 6 unsaturated fatty acids, which are involved in the maintenance of membrane structure and function, the regulation of cholesterol synthesis and transport, and the prevention

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of water loss from the skin. Delta 6 unsaturated fatty acids also serve as precursors of the eicosanoids, including the prostaglandins and leukotrienes (Horrobin, 1992, Prog. Lipid Res. 31:163-194). The double bond at the 6 position of delta 6 unsaturated fatty acids is introduced by a class of enzymes known as delta 6 desaturases.

Deficiencies in linoleic acid and ALA derivatives have been associated with skin diseases, diabetic complications, inflammatory and autoimmune disorders, cardiovascular disorders, complications of viral infection, and retinal dysfunction. For example, a deficiency in gamma-linolenic acid (GLA), which is produced from linoleic acid by the action of the enzyme delta 6 desaturase, can arise from the decreased activity of this enzyme that occurs in aging, stress, diabetes, eczema, and some infections, or from increased catabolism of GLA due to oxidation or rapid cell division, as occurs in inflammation or cancer. Clinical trials have demonstrated that dietary GLA supplementation can be effective in treating a number of conditions that are associated with GLA deficiency, e.g., atopic eczema, mastalgia, diabetic neuropathy, viral infections, and some forms of cancer (Horrobin, 1990, Rev. Contemp. Pharmacother. 1:1-45).

Delta 6 desaturase is an example of a fatty acid desaturase. Fatty acid desaturases are enzymes that introduce a double bond into the carbon chain of fatty acids. They play vital roles in the biosynthesis of polyunsaturated fatty acids, including the essential fatty acids. Fatty acid desaturases are present in soluble and membrane-associated forms and require electron donors (for example, cytochrome b5) for their functioning.

Delta 6 desaturases catalyze the rate-limiting steps in the biosyntheses of the linoleic and ALA group EFAs shown in Figure 1. End products of the linoleic acid pathway include the eicosanoids (prostaglandins and leukotrienes). The end product of the ALA pathway is docosahexaenoic acid (DHA), an important component of membranes in the vertebrate retina. DHA is highly specific for retina and represents more than 50% of the fatty acids in the rod outer segment (ROS). It appears that DHA is important in maintaining the normal structure and function of the retina (Anderson et al., 1992, Neurobiology of Essential Fatty Acids, Bazan et al., eds., Plenum Press, New York, pages 285-294). Increased dietary consumption of DHA and its precursor, eicosapentaenoic acid, from seal meat and fish has been

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linked to an increased incidence of macular degeneration in Greenland Eskimos (Rosenberg, 1987, Arct. Med. Res. 46:64-70).

Certain delta 6 desaturases have been cloned from plants. For example, a delta 6 desaturase has been cloned from borage (Sayanova et al., 1997, Proc. Natl. Acad. Sci. USA 94:4211-4216). This delta 6 desaturase is unusual in that its cytochrome b5 electron donor is present as an N-terminal extension of the enzyme rather than being synthesized as a separate protein. The borage delta 6 desaturase has been shown to be functional, in that transfer of the cloned gene encoding it to tobacco results in the synthesis of high levels of GLA and octadecatetraenoic acid (OTA) in the transgenic tobacco leaves. GLA and OTA are the products of delta 6 desaturase activity on linoleic acid and ALA, respectively.

Based on its hydropathy profile, the borage delta 6 desaturase appears to be a membrane-bound protein. Examination of the amino acid sequence of the borage enzyme, as well as the amino acid sequences of membrane-bound desaturases from a wide variety of organisms, has revealed three regions of conserved short motifs containing histidine residues (HX(3 or 4)H, HX(2 or 3)HH, and HX(2 or 3)HH) having a conserved spacing from each other (Shanklin et al., Biochemistry, 1994, 33:12787-12794).

A DNA sequence has been isolated from sunflower embryos that, judging from its sequence, appears to encode a delta 6 desaturase having a cytochrome b5-like moiety fused to its N-terminus (Sperling et al., 1995, Eur. J. Biochem. 232:798-805).

SUMMARY OF THE INVENTION

The present invention is directed to novel human DNA sequences that encode a delta 6 fatty acid desaturase, cytochrome b5-related protein (CYB5RP). The present invention includes genomic CYB5RP DNA as well as cDNA that encodes the CYB5RP protein. The genomic CYB5RP DNA is substantially free from other nucleic acids and has the nucleotide sequence shown in SEQ.ID.NO.:1. The cDNA encoding CYB5RP protein is substantially free from other nucleic acids and has the nucleotide sequence shown in SEQ.ID.NO.:2. Also provided is CYB5RP protein encoded by the novel DNA sequences. The CYB5RP protein is substantially free from other proteins and has the amino acid sequence shown in SEQ.ID.NO.:3.

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(SEQ.ID.NO.:3).

Methods of expressing CYB5RP protein in recombinant systems are provided. Also provided are methods of producing delta 6 unsaturated fatty acids using DNA encoding CYB5RP or using CYB5RP protein.

5 BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 depicts the enzymatic conversions involved in the linoleic acid (n-3) and alpha-linolenic acid (n-6) pathways of essential fatty acid synthesis.

Figure 2A-G shows the genomic DNA sequence of the CYB5RP gene (SEQ.ID.NO.:1). Underlined nucleotides in capitals represent exons. The start ATG codon at position 544 in exon 1 and the stop TGA codon at position 18,103 in exon 12 are shown in bold. The putative polyadenylation signal ATTAAA located approximately 20 base pairs upstream of the polyA tail is shown in bold italics (position 18,373 in exon 12). DNA sequence upstream of exon 1 represents a putative promoter region of the CYB5RP gene., as indicated by the presence of the TATA box at position 353 (underlined bold)..

Figure 3A-C shows the cDNA sequence (SEQ.ID.NO.:2) and the amino acid sequence (SEQ.ID.NO.:3) of CYB5RP. The region encompassing amino acids 1-102 represents the cytochrome b5 domain. The region encompassing amino acids 182-186 represents HIS BOX 1. The region encompassing amino acids 219-223 represents HIS BOX 2. The region encompassing amino acids 383-387 represents HIS BOX 3.

Figure 4 shows a portion of the cDNA sequence (SEQ.ID.NO.:4) and a portion of the amino acid sequence (SEQ.ID.NO.:5) of mouse CYB5RP.

Figure 5A shows a Kyte-Doolittle hydropathy plot of CYB5RP. Figure 5B shows the proposed membrane topology of CYB5RP based on its hydropathy plot. This membrane topology is similar to that proposed for other membrane-bound fatty acid desaturases (Shanklin et al., Biochemistry, 1994, 33:12787-12794). The amino acids shown in Figure 5B are portions of

Figure 6 shows the output of the Profilescan program from the Wisconsin GCG package. The upper amino acid sequence is from CYB5RP (positions 31-78 of SEQ. ID. NO.3). The lower amino acid sequence is positions 1-48 of the cytochrome b5 profile (SEQ. ID. NO.:6.). The output shows that CYB5RP

contains a profile typical for the heme-binding domain of the cytochrome b5 protein family. Importantly, the region of identity includes the invariant HPGG motif, where histidine represents a heme axial ligand for iron.

Figure 7A and B show the results of BlastP searches of the GenBank database using the full-length CYB5RP amino acid sequence as the query. Figure 7A 5 shows the hit with highest homology, a hypothetical protein from sunflower. The sunflower protein and CYB5RP share three His boxes (boxed) in which the spacing between the His boxes is conserved. Also boxed is the HPGG motif typical for the heme-binding domain of the cytochrome b5 protein family. In both proteins the first histidine of the third His box is replaced by glutamine (a typical feature of desaturases 10 with delta 6 specificity). The upper amino acid sequences shown are from CYB5RP and are portions of SEQ. ID. NO.3. The lower amino acid sequences shown are portions of the amino acid sequence of the hypothetical protein from sunflower (Sperling et al., 1995, Eur. J. Biochem. 232:798-805). The sequence shown as positions 348-432 is SEQ. ID. NO.:7. The sequence shown as positions 22-74 is 15 SEQ. ID. NO.:8. The sequence shown as positions 152-227 is SEQ. ID. NO.:9. Figure 7B shows the hit with the second highest homology, a delta 6 desaturase from Borago oficinalis (Sayanova et al., 1997, Proc. Natl. Acad. Sci. USA 94:4211-4216). The Borago protein and CYB5RP also share three His boxes with conserved spacing, as well as the HPGG motif. In both proteins the first histidine of the third His box is 20 replaced by glutamine (a typical feature of desaturases with delta 6 specificity). The upper amino acid sequences shown are from CYB5RP and are portions of SEQ. ID. NO.3. The lower amino acid sequences shown are portions of the amino acid sequence of the Borago delta 6 desaturase. The sequence shown as positions 338-424 is SEQ. ID. NO.:10. The sequence shown as positions 12-64 is SEQ. ID. NO.:11. 25 The sequence shown as positions 153-220 is SEQ. ID. NO.:12.

Figure 8 shows additional results of BlastP searches of the GenBank database using the CYB5RP protein as the query. Figure 8 shows the amino acid alignment between the CYB5RP protein and a delta 6 desaturase from Synechocystis sp. (strain pcc 6803) performed by the BlastP program. The Synechocystis delta 6 desaturase and CYB5RP share three His boxes, two of which are shown in Figure 8 (boxed). In both proteins the first histidine of the third His box is replaced by glutamine (a typical feature of desaturases with delta 6 specificity). The CYB5RP

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sequence shown is a portion of SEQ. ID. NO.3. The *Synechocystis* sequence shown is SEQ. ID. NO:13.

Figure 9A shows the expression pattern of the CYB5RP gene in 9 human tissues, as determined by RT-PCR amplification with 21 cycles. Expression is detected in human retina, kidney, pancreas, placenta, and brain. Figure 9B shows the results of the analogous experiments performed with 25 cycles of amplification. Expression of the CYB5RP gene is seen in all the human tissues studied.

DETAILED DESCRIPTION OF THE INVENTION

For the purposes of this invention:

"Substantially free from other proteins" means at least 90%, preferably 95%, more preferably 99%, and even more preferably 99.9%, free of other proteins. Thus, a CYB5RP protein preparation that is substantially free from other proteins will contain, as a percent of its total protein, no more than 10%, preferably no more than 5%, more preferably no more than 1%, and even more preferably no more than 0.1%, of non-CYB5RP proteins. Whether a given CYB5RP protein preparation is substantially free from other proteins can be determined by such conventional techniques of assessing protein purity as, e.g., sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) combined with appropriate detection methods, e.g., silver staining or immunoblotting.

"Substantially free from other nucleic acids" means at least 90%, preferably 95%, more preferably 99%, and even more preferably 99.9%, free of other nucleic acids. Thus, a CYB5RP DNA preparation that is substantially free from other nucleic acids will contain, as a percent of its total nucleic acid, no more than 10%, preferably no more than 5%, more preferably no more than 1%, and even more preferably no more than 0.1%, of non-CYB5RP nucleic acids. Whether a given CYB5RP DNA preparation is substantially free from other nucleic acids can be determined by such conventional techniques of assessing nucleic acid purity as, e.g., agarose gel electrophoresis combined with appropriate staining methods, e.g., ethidium bromide staining, or by sequencing.

"Substantially the same biological activity as CYB5RP" means being able to introduce a double bond into the 6 position of linoleic acid under conditions in which CYB5RP is able to introduce a double bond into the 6 position of linoleic acid.

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A "conservative amino acid substitution" refers to the replacement of one amino acid residue by another, chemically similar, amino acid residue. Examples of such conservative substitutions are: substitution of one hydrophobic residue (isoleucine, leucine, valine, or methionine) for another; substitution of one polar residue for another polar residue of the same charge (e.g., arginine for lysine; glutamic acid for aspartic acid); substitution of one aromatic amino acid (tryptophan, tyrosine, or phenylalanine) for another.

The present invention relates to the identification and cloning of cytochrome b5-related protein (CYB5RP), a gene which encodes a human delta 6 fatty acid desaturase. The gene is present on PAC clones 759J12, 756B3, 519O13, and 466A11 from an area of human chromosome 11q12 that has been shown to contain a gene related to Best's macular dystrophy (Cooper et al., 1997, Genomics 41:185-192; Stöhr et al., 1997, Genome Res. 8:48-56; Graff et al., 1997, Hum. Genet. 101: 263-279). This linkage between the chromosomal location of the CYB5RP gene and the location of the gene related to Best's macular dystrophy can be used diagnostically by identifying restriction fragment length polymorphisms (RFLPs) in the vicinity of the CYB5RP gene, e.g., in SEQ.ID.NO.:1. Such RFLPs will be associated with the Best's macular dystrophy gene and thus can be used to identify individuals carrying disease-causing forms of the Best's macular dystrophy gene.

CYB5RP was identified as an EST hit in sequence scanning data from PAC clones from human chromosome 11q12. In addition, a full length cDNA of CYB5RP was recovered from a human retina cDNA library. The genomic region of CYB5RP has been sequenced and the exon/intron organization of CYB5RP has been determined. The CYB5RP gene has 12 exons. The promoter region of CYB5RP was identified upstream of the 5' UTR by detecting consensus elements required for eukaryotic transcription. The expression pattern of CYB5RP was determined by RT-PCR analysis in 9 human tissues. The CYB5RP gene is expressed predominantly in human retina, kidney, pancreas, and placenta; lower levels of expression are also detected in brain, heart, lung, liver, and skeletal muscle. Bioinformatic analysis revealed significant homology to a group of plant and bacterial fatty acid desaturases. All of the typical amino acid motifs present in these fatty acid desaturases are also present in CYB5RP. Kyte-Doolittle algorithm analysis predicts a transmembrane organization typical of fatty acid desaturases for CYB5RP (see Figure 5). CYB5RP is

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unusual in that it contains a cytochrome b5 region in its N terminus. While many fatty acid desaturases utilize cytochrome b5 as an electron donor, most have not incorporated this cytochrome as part of their polypeptide chain.

That CYB5RP is a fatty acid desaturase is shown by the following evidence:

- (1) CYB5RP possesses significant homology to a group of plant and microbial fatty acid desaturases;
- (2) Like other fatty acid desaturases, CYB5RP has three conserved histidine boxes, with correct spacing between the boxes; and
- (3) The predicted membrane topology of CYB5RP is similar to that of known fatty acid desaturases.

That CYB5RP is a delta 6 fatty acid desaturase is shown by the following evidence:

- (1) CYB5RP contains a cytochrome b5-like moiety fused to its N-terminus. The only two fatty acid desaturases that contain cytochrome b5-like moiety fused to their N-termini are known or suspected to be delta 6 desaturases.
- (2) The only two plant desaturases that are known or suspected to introduce a double bond in the 6 position have an atypical His box 3 (QI/LEHH), with a Q in the first position rather than an H. CYB5RP has the same atypical His Box 3.
- (3) The only bacterial desaturase that is known to introduce a double bond in the 6 position has an atypical His box 3 (QVTHH), with a Q in the first position rather than an H. CYB5RP has the same atypical His Box 3.

CYB5RP is a target for the development of drugs for the treatment of disorders of lipid metabolism and for the treatment of conditions that require the modulation of the biosynthesis of prostaglandins and leukotrienes (asthma, pain, etc.). CYB5RP is also a target for the development of drugs for use in treating skin diseases, diabetic complications, reproductive disorders, including breast pain and premenstrual syndrome, inflammatory and autoimmune disorders, cardiovascular disorders, complications of viral infections, and various forms of retinal degeneration, including age-related macular degeneration.

CYB5RP is homologous to a delta 6 desaturase from *Borago oficinalis* (see Figure 7B). Both CYB5RP and this *Borago* delta 6 desaturase, unlike desaturases from higher plants, are unusual in containing a cytochrome b5-like

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domain fused to their N-termini (Sayanova et al., 1997, Proc. Natl. Acad. Sci. USA 94:4211-4216; hereinafter "Sayanova"). The *Borago* desaturase has been expressed in transgenic tobacco, resulting in high levels of delta 6 desaturated fatty acids in the transgenic tobacco leaves, including high levels of γ-linolenic acid (GLA) (Sayanova). Given the medical importance of GLA, Sayanova proposed that transgenic plants, expressing the *Borago* delta 6 desaturase, would be valuable as sources of GLA. Similarly, CYB5RP, expressed in transgenic plants, is expected to provide a valuable source of GLA.

The present invention provides DNA encoding CYB5RP that is substantially free from other nucleic acids. The present invention also provides recombinant DNA molecules encoding CYB5RP. The present invention provides DNA molecules substantially free from other nucleic acids comprising the nucleotide sequence shown in Figure 2 as SEQ.ID.NO.:1. Analysis of SEQ.ID.NO.:1 revealed that this genomic sequence defines a gene having 12 exons. These exons collectively have an open reading frame that encodes a protein of 445 amino acids. When an alternatively spliced exon 8 is used, a CYB5RP protein of 433 amino acids, lacking amino acids 317-328, is produced. Thus, the present invention includes two cDNA molecules, encoding two forms of CYB5RP protein, that are substantially free from other nucleic acids. The first cDNA is shown in Figure 3 and has the nucleotide sequence SEQ.ID.NO.:2. The second cDNA is identical to the first, except that it does not contain the nucleotides at positions 1,019-1,054.

The present invention includes DNA molecules substantially free from other nucleic acids comprising the coding region of SEQ.ID.NO.:2. Accordingly, the present invention includes DNA molecules substantially free from other nucleic acids having a sequence comprising positions 71-1,405 of SEQ.ID.NO.:2. The present invention also includes DNA molecules substantially free from other nucleic acids having a sequence comprising positions 71-1,405 of SEQ.ID.NO.:2, except that the nucleotides at positions 1,019-1,054 are missing. Also included in the present invention are recombinant DNA molecules having a nucleotide sequence comprising positions 71-1,405 of SEQ.ID.NO.:2 and recombinant DNA molecules having a nucleotide sequence comprising positions 71-1,405 of SEQ.ID.NO.:2 with the exception that positions 1,019-1,054 are missing.

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The novel DNA sequences of the present invention encoding CYB5RP, in whole or in part, can be linked with other DNA sequences, *i.e.*, DNA sequences to which CYB5RP is not naturally linked, to form "recombinant DNA molecules" encoding CYB5RP. Such other sequences can include DNA sequences that control transcription or translation such as, *e.g.*, translation initiation sequences, promoters for RNA polymerase II, transcription or translation termination sequences, enhancer sequences, sequences that control replication in microorganisms, sequences that confer antibiotic resistance, or sequences that encode a polypeptide "tag" such as, *e.g.*, a polyhistidine tract or the myc epitope. The novel DNA sequences of the present invention can be inserted into vectors such as plasmids, cosmids, viral vectors, P1 artificial chromosomes, or yeast artificial chromosomes.

Included in the present invention are DNA sequences that hybridize to at least one of SEQ.ID.NOs.:1 or 2 under stringent conditions. By way of example, and not limitation, a procedure using conditions of high stringency is as follows: Prehybridization of filters containing DNA is carried out for 2 hr. to overnight at 65°C in buffer composed of 6X SSC, 5X Denhardt's solution, and 100 µg/ml denatured salmon sperm DNA. Filters are hybridized for 12 to 48 hrs at 65°C in prehybridization mixture containing 100 µg/ml denatured salmon sperm DNA and 5-20 X 10⁶ cpm of ³²P-labeled probe. Washing of filters is done at 37°C for 1 hr in a solution containing 2X SSC, 0.1% SDS. This is followed by a wash in 0.1X SSC, 0.1% SDS at 50°C for 45 min. before autoradiography.

Other procedures using conditions of high stringency would include either a hybridization carried out in 5XSSC, 5X Denhardt's solution, 50% formamide at 42°C for 12 to 48 hours or a washing step carried out in 0.2X SSPE, 0.2% SDS at 65°C for 30 to 60 minutes.

Reagents mentioned in the foregoing procedures for carrying out high stringency hybridization are well known in the art. Details of the composition of these reagents can be found in, e.g., Sambrook, Fritsch, and Maniatis, 1989, Molecular Cloning: A Laboratory Manual, second edition, Cold Spring Harbor Laboratory Press. In addition to the foregoing, other conditions of high stringency which may be used are well known in the art.

The degeneracy of the genetic code is such that, for all but two amino acids, more than a single codon encodes a particular amino acid. This allows for the

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construction of synthetic DNA that encodes the CYB5RP protein where the nucleotide sequence of the synthetic DNA differs significantly from the nucleotide sequence of SEQ.ID.NO.:2, but still encodes the same CYB5RP protein shown as SEQ.ID.NO.:3. Such synthetic DNAs are intended to be within the scope of the present invention. Also with the scope of the present invention are synthetic DNAs that encode a CYB5RP protein lacking amino acids 317-328 of SEQ.ID.NO.:3.

Another aspect of the present invention includes host cells that have been engineered to contain and/or express DNA sequences encoding CYB5RP protein. Such recombinant host cells can be cultured under suitable conditions to produce CYB5RP protein. An expression vector containing DNA encoding CYB5RP protein can be used for expression of CYB5RP protein in a recombinant host cell. Recombinant host cells may be prokaryotic or eukaryotic, including but not limited to, bacteria such as E. coli, fungal cells such as yeast, mammalian cells including, but not limited to, cell lines of human, bovine, porcine, monkey and rodent origin, plant cells such as tobacco, and insect cells including but not limited to Drosophila and silkworm derived cell lines. Cell lines derived from mammalian species which are suitable for recombinant expression of CYB5RP protein and which are commercially available, include but are not limited to, L cells L-M(TK-) (ATCC CCL 1.3), L cells L-M (ATCC CCL 1.2), 293 (ATCC CRL 1573), Raji (ATCC CCL 86), CV-1 (ATCC CCL 70), COS-1 (ATCC CRL 1650), COS-7 (ATCC CRL 1651), CHO-K1 (ATCC CCL 61), 3T3 (ATCC CCL 92), NIH/3T3 (ATCC CRL 1658), HeLa (ATCC CCL 2), C127I (ATCC CRL 1616), BS-C-1 (ATCC CCL 26) and MRC-5 (ATCC CCL 171).

A variety of mammalian expression vectors can be used to express recombinant CYB5RP in mammalian cells. Commercially available mammalian expression vectors which are suitable include, but are not limited to, pMC1neo (Stratagene), pSG5 (Stratagene), pcDNAI and pcDNAIamp, pcDNA3, pcDNA3.1, pCR3.1 (Invitrogen), EBO-pSV2-neo (ATCC 37593), pBPV-1(8-2) (ATCC 37110), pdBPV-MMTneo(342-12) (ATCC 37224), pRSVgpt (ATCC 37199), pRSVneo (ATCC 37198), and pSV2-dhfr (ATCC 37146). Following expression in recombinant cells, CYB5RP can be purified by conventional techniques to a level that is substantially free from other proteins. A description of vectors that can be used to express CYB5RP can be found in, e.g., Goeddel, ed., 1990, Meth. Enzymol. vol. 185 or Perbal, 1988, A Practical Guide to Molecular Cloning, John Wiley and Sons, Inc.

The present invention includes CYB5RP protein substantially free from other proteins. The amino acid sequence of the full-length CYB5RP protein is shown in Figure 3 as SEQ.ID.NO.:3. Thus, the present invention includes CYB5RP protein substantially free from other proteins having the amino acid sequence SEQ.ID.NO.:3. Also included in the present invention is a CYB5RP protein that is produced from an alternatively spliced CYB5RP mRNA where the protein has the amino acid sequence of SEQ.ID.NO.:3 with the exception that amino acids 317-328 are missing.

As with many proteins, it is possible to modify many of the amino acids of CYB5RP and still retain substantially the same biological activity as the 10 original protein. Thus, the present invention includes modified CYB5RP proteins which have amino acid deletions, additions, or substitutions but that still retain substantially the same biological activity as CYB5RP. It is generally accepted that single amino acid substitutions do not usually alter the biological activity of a protein (see, e.g., Molecular Biology of the Gene, Watson et al., 1987, Fourth Ed., The 15 Benjamin/Cummings Publishing Co., Inc., page 226; and Cunningham & Wells, 1989, Science 244:1081-1085). Accordingly, the present invention includes polypeptides where one amino acid substitution has been made in SEQ.ID.NO.:3 wherein the polypeptides still retain substantially the same biological activity as CYB5RP. The present invention also includes polypeptides where two or more 20 amino acid substitutions have been made in SEQ.ID.NO.:3 wherein the polypeptides still retain substantially the same biological activity as CYB5RP. In particular, the present invention includes embodiments where the above-described substitutions are conservative substitutions. In particular, the present invention includes embodiments where the above-described substitutions do not occur in the His boxes of CYB5RP. 25 In particular, the present invention includes embodiments where the above-described substitutions do not occur in positions where the amino acid present in those positions in CYB5RP is the same as the amino acid present in the corresponding position of the sunflower protein depicted in Figure 1 of Sperling et al., 1995, Eur. J. Biochem. 232:798-805 when these two proteins are aligned by BLASTP analysis. In particular, 30 the present invention includes embodiments where the above-described substitutions do not occur in positions where the amino acid present in those positions in CYB5RP is the same as the amino acid present in the corresponding position of the

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CCCTCTACCCCTGTCCCATCAGGC (SEQ.ID.NO.:15)

One of skill in the art would recognize that many other primer pairs based upon SEQ.ID.NO.:2 would also be suitable.

PCR reactions can be carried out with a variety of thermostable enzymes including but not limited to AmpliTaq, AmpliTaq Gold, or Vent polymerase. For AmpliTaq, reactions can be carried out in 10 mM Tris-Cl, pH 8.3, 2.0 mM MgCl₂, 200 μM for each dNTP, 50 mM KCl, 0.2 μM for each primer, 10 ng of DNA template, 0.05 units/μl of AmpliTaq. The reactions are heated at 95°C for 3 minutes and then cycled 35 times using the cycling parameters of 95°C, 20 seconds, 62°C, 20 seconds, 72°C, 3 minutes. In addition to these conditions, a variety of suitable PCR protocols can be found in PCR Primer, A Laboratory Manual, edited by C.W. Dieffenbach and G.S. Dveksler, 1995, Cold Spring Harbor Laboratory Press; or PCR Protocols: A Guide to Methods and Applications, Michael et al., eds., 1990, Academic Press.

A suitable cDNA library from which a clone encoding CYB5RP can be isolated would be Human Retina 5'-stretch cDNA library in lambda gt10 or lambda gt11 vectors (catalog numbers HL1143a and HL1132b, Clontech, Palo Alto, CA). The primary clones of such a library can be subdivided into pools with each pool containing approximately 20,000 clones and each pool can be amplified separately.

By this method, a cDNA fragment encoding an open reading frame of either 445 amino acids (SEQ.ID.NO.:3) or an open reading frame of 433 amino acids (SEQ.ID.NO.:3 lacking the amino acids at positions 317-328) can be obtained. This cDNA fragment can be cloned into a suitable cloning vector or expression vector. For example, the fragment can be cloned into the mammalian expression vector pcDNA3.1 (Invitrogen, San Diego, CA). CYB5RP protein can then be produced by transferring an expression vector encoding CYB5RP or portions thereof into a suitable host cell and growing the host cell under appropriate conditions. CYB5RP protein can then be isolated by methods well known in the art.

As an alternative to the above-described PCR method, a cDNA clone encoding CYB5RP can be isolated from a cDNA library using as a probe oligonucleotides specific for CYB5RP and methods well known in the art for screening cDNA libraries with oligonucleotide probes. Such methods are described

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in, e.g., Sambrook et al., 1989, Molecular Cloning: A Laboratory Manual; Cold Spring Harbor Laboratory, Cold Spring Harbor, New York; Glover, D.M. (ed.), 1985, DNA Cloning: A Practical Approach, MRL Press, Ltd., Oxford, U.K., Vol. I, II. Oligonucleotides that are specific for CYB5RP and that can be used to screen cDNA libraries can be readily designed based upon the cDNA sequence of CYB5RP shown

in SEQ.ID.NO.:2 and can be synthesized by methods well-known in the art. Genomic clones containing the CYB5RP gene can be obtained from commercially available human PAC or BAC libraries available from Research Genetics, Huntsville, AL. PAC clones containing the CYB5RP gene (e.g., PAC clones 759J12, 756B3, 519O13, and 466A11) are commercially available from Research Genetics, Huntsville, AL (Catalog number for individual PAC clones is RPCI.C). Alternatively, one may prepare genomic libraries, especially in P1 artificial chromosome vectors, from which genomic clones containing the CYB5RP can be isolated, using probes based upon the CYB5RP sequences disclosed herein. Methods of preparing such libraries are known in the art (Ioannou et al., 1994, Nature Genet. 6:84-89).

The present invention also provides oligonucleotide probes, based upon SEQ.ID.NO.:2 that can be used to determine the level of CYB5RP RNA in a sample. In particular, the present invention includes DNA oligonucleotides comprising at least 18 contiguous nucleotides of SEQ.ID.NO.:2. Also provided by the present invention are corresponding RNA oligonucleotides. The DNA or RNA oligonucleotide probes can be packaged in kits.

In addition to the utilities described above, the present invention makes possible the recombinant expression of the CYB5RP protein in various cell types. In particular, it is advantageous to recombinantly express CYB5RP in plant cells. Such expression in plant cells provides a method for the production of high levels of valuable EFAs such as GLA and OTA in the recombinant plant cells. An example of such recombinant expression of a delta 6 fatty acid desaturase, in that case from borage, is described in Sayanova et al., 1997, Proc. Natl. Acad. Sci. USA 94:4211-4216 (Sayanova). The recombinant expression of the borage delta 6 desaturase led to the production of high levels of GLA and OTA in the leaves of the tobacco plants in which it was expressed. The procedures described in Sayanova can be easily adapted to express CYB5RP in tobacco, thus providing an additional, useful way to produce

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large amounts of valuable EFAs. Known methods of recombinantly expressing genes in other plant species beside tobacco can be used to express CYB5RP in those other species.

The present invention also makes possible the development of assays which measure the biological activity of the CYB5RP protein. Such assays using recombinantly expressed CYB5RP protein are especially of interest.

Assays for CYB5RP protein activity can be used to screen libraries of compounds or other sources of compounds to identify compounds that are activators or inhibitors of the activity of CYB5RP protein. Such identified compounds can serve as "leads" for the development of pharmaceuticals that can be used to modulate the activity of CYB5RP in patients suffering from conditions where that activity is abnormal, e.g., skin diseases, diabetic complications, inflammatory and autoimmune disorders, cardiovascular disorders, complications of viral infection, and retinal dysfunction such as macular degeneration.

Such assays may comprise:

- (a) recombinantly expressing CYB5RP protein in a host cell;
- (b) measuring the biological activity of the recombinantly expressed CYB5RP protein in the presence and in the absence of a substance suspected of being an activator or an inhibitor of CYB5RP protein;

where a change in the biological activity of the recombinantly expressed CYB5RP protein in the presence as compared to the absence of the substance indicates that the substance is an activator or an inhibitor of CYB5RP protein.

In particular embodiments, the biological activity of the recombinantly expressed CYB5RP protein is the ability to introduce a double bond into the 6 position of linoleic acid or alpha-linoleic acid.

In some embodiments, it may be advantageous to insert additional steps between steps (a) and (b). Such additional steps might include lysing the host cell and fractionating its contents in order to partially purify the recombinantly expressed CYB5RP, thus facilitating exposure of the recombinantly expressed CYB5RP to the substance as well as to any substrate used in the assay.

The present invention includes activators and inhibitors identified by the methods described herein as well as pharmaceutical compositions comprising

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such activators and inhibitors. The activators and inhibitors are generally combined with pharmaceutically acceptable carriers before use to form pharmaceutical compositions. Examples of such carriers and methods of formulation of pharmaceutical compositions containing activators or inhibitors and carriers can be found in Remington's Pharmaceutical Sciences. To form a pharmaceutically acceptable composition suitable for effective administration, such compositions will contain an effective amount of the activator or inhibitor.

Therapeutic or prophylactic compositions are administered to an individual in amounts sufficient to treat or prevent conditions where CYB5RP activity is abnormal. The effective amount can vary according to a variety of factors such as the individual's condition, weight, sex and age. Other factors include the mode of administration. The appropriate amount can be determined by a skilled physician.

Compositions can be used alone at appropriate dosages. Alternatively, co-administration or sequential administration of other agents can be desirable.

The compositions can be administered in a wide variety of therapeutic dosage forms in conventional vehicles for administration. For example, the compositions can be administered in such oral dosage forms as tablets, capsules (each including timed release and sustained release formulations), pills, powders, granules, elixirs, tinctures, solutions, suspensions, syrups and emulsions, or by injection. Likewise, they can also be administered in intravenous (both bolus and infusion), intraperitoneal, subcutaneous, topical with or without occlusion, or intramuscular form, all using forms well known to those of ordinary skill in the pharmaceutical arts.

Advantageously, compositions can be administered in a single daily dose, or the total daily dosage can be administered in divided doses of two, three or four times daily. Furthermore, compositions can be administered in intranasal form via topical use of suitable intranasal vehicles, or via transdermal routes, using those forms of transdermal skin patches well known to those of ordinary skill in that art. To be administered in the form of a transdermal delivery system, the dosage administration will, of course, be continuous rather than intermittent throughout the dosage regimen.

The dosage regimen utilizing the compositions is selected in accordance with a variety of factors including type, species, age, weight, sex and medical condition of the patient; the severity of the condition to be treated; the route

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of administration; the renal, hepatic and cardiovascular function of the patient; and the particular composition thereof employed. A physician or veterinarian of ordinary skill can readily determine and prescribe the effective amount of the composition required to prevent, counter or arrest the progress of the condition. Optimal precision in achieving concentrations of composition within the range that yields efficacy without toxicity requires a regimen based on the kinetics of the composition's availability to target sites. This involves a consideration of the distribution, equilibrium, and elimination of a composition.

The present invention also includes antibodies to the CYB5RP protein. Such antibodies may be polyclonal antibodies or monoclonal antibodies. The antibodies of the present invention are raised against the entire CYB5RP protein or against suitable antigenic fragments of the protein that are coupled to suitable carriers, e.g., serum albumin or keyhole limpet hemocyanin, by methods well known in the art. Methods of identifying suitable antigenic fragments of a protein are known in the art. See, e.g., Hopp & Woods, 1981, Proc. Natl. Acad. Sci. USA 78:3824-3828; and Jameson & Wolf, 1988, CABIOS (Computer Applications in the Biosciences) 4:181-186.

For the production of polyclonal antibodies, CYB5RP protein or an antigenic fragment, coupled to a suitable carrier, is injected on a periodic basis into an appropriate non-human host animal such as, e.g., rabbits, sheep, goats, rats, mice. The animals are bled periodically and sera obtained are tested for the presence of antibodies to the injected antigen. The injections can be intramuscular, intraperitoneal, subcutaneous, and the like, and can be accompanied with adjuvant.

For the production of monoclonal antibodies, CYB5RP protein or an antigenic fragment, coupled to a suitable carrier, is injected into an appropriate non-human host animal as above for the production of polyclonal antibodies. In the case of monoclonal antibodies, the animal is generally a mouse. The animal's spleen cells are then immortalized, often by fusion with a myeloma cell, as described in Kohler & Milstein, 1975, Nature 256:495-497. For a fuller description of the production of monoclonal antibodies, see Antibodies: A Laboratory Manual, Harlow & Lane, eds., Cold Spring Harbor Laboratory Press, 1988.

Gene therapy may be used to introduce CYB5RP polypeptides into the cells of target organs, e.g., the pigmented epithelium of the retina or other parts of the

retina. Nucleotides encoding CYB5RP polypeptides can be ligated into viral vectors which mediate transfer of the nucleotides by infection of recipient cells. Suitable viral vectors include retrovirus, adenovirus, adeno-associated virus, herpes virus, vaccinia virus, and polio virus based vectors. Alternatively, nucleotides encoding CYB5RP polypeptides can be transferred into cells for gene therapy by non-viral techniques including receptor-mediated targeted transfer using ligand-nucleotide conjugates, lipofection, membrane fusion, or direct microinjection. These procedures and variations thereof are suitable for ex vivo as well as in vivo gene therapy. Gene therapy with CYB5RP polypeptides will be particularly useful for the treatment of diseases where it is beneficial to elevate CYB5RP activity.

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The present invention is not to be limited in scope by the specific embodiments described herein. Indeed, various modifications of the invention in addition to those described herein will become apparent to those skilled in the art from the foregoing description. Such modifications are intended to fall within the scope of the appended claims.

Various publications are cited herein, the disclosures of which are incorporated by reference in their entireties.

WO 00/21557 PCT/US99/23253

WHAT IS CLAIMED:

1. A recombinant DNA molecule encoding a polypeptide having the amino acid sequence of SEQ.ID.NO.:3.

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2. A recombinant DNA molecule comprising a nucleotide sequence selected from the group consisting of:

SEQ.ID.NO.:1;

SEQ.ID.NO.:2;

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SEQ.ID.NO.:2 lacking positions 1,019-1,054; positions 71-1,405 of SEQ.ID.NO.:2; and

positions 71-1,405 of SEQ.ID.NO.:2 lacking positions 1,019-1,054.

- 3. A DNA molecule that hybridizes under stringent conditions to the DNA molecule of claim 2.
 - 4. An expression vector comprising the DNA of claim 1.

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- 5. A recombinant host cell comprising the DNA of claim 1.
- 6. A CYB5RP protein, substantially free from other proteins, having an amino acid sequence selected from the group consisting of SEQ.ID.NO.:3 and SEQ.ID.NO.:3 lacking positions 317-328.

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- 7. The CYB5RP protein of claim 6 containing a single amino acid substitution.
- 8. The CYB5RP protein of claim 7 where the substitution is a conservative substitution.
 - 9. The CYB5RP protein of claim 6 containing amino acid substitutions where the substitutions do not occur in positions where the amino acid

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present in CYB5RP at those positions is also present in the corresponding position in the delta 6 desaturase from sunflower when CYB5RP and the delta 6 desaturase from sunflower are aligned by BLASTP analysis or where the substitutions do not occur in positions where the amino acid present in CYB5RP at those positions is also present in the corresponding position in the delta 6 desaturase from *Synechocystis* when CYB5RP and the delta 6 desaturase from *Synechocystis* are aligned by BLASTP analysis or where the substitutions do not occur in positions where the amino acid present in CYB5RP at those positions is also present in the corresponding position in the delta 6 desaturase from borage when CYB5RP and the delta 6 desaturase from borage are aligned by BLASTP analysis.

- 10. An antibody that binds specifically to the CYB5RP protein of claim 6.
- 15 11. A DNA or RNA oligonucleotide probe comprising at least 18 contiguous nucleotides of at least one of the sequences of claim 2.
 - 12. A method for determining whether a substance is an activator or an inhibitor of CYB5RP protein comprising:
 - (a) recombinantly expressing the CYB5RP protein of claim 6 in a host cell;
 - (b) measuring the biological activity of the recombinantly expressed CYB5RP protein in the presence and in the absence of a substance suspected of being an activator or an inhibitor of CYB5RP protein;
 - where a change in the biological activity of the recombinantly expressed CYB5RP protein in the presence as compared to the absence of the substance indicates that the substance is an activator or an inhibitor of CYB5RP protein.
- The method of claim 12 where the biological activity of CYB5RP protein is the ability to introduce a double bond into the 6 position of linoleic acid.

- 14. A pharmaceutical composition comprising an activator or an inhibitor of CYB5RP.
- 15. A method of treating macular degeneration comprising
 administering to a patient an effective amount of the pharmaceutical composition of claim 14.

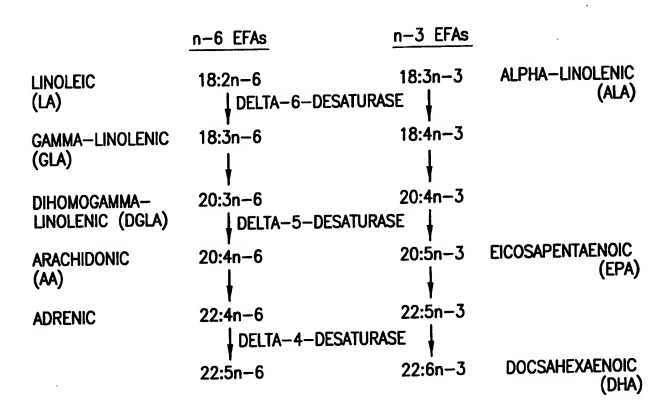


FIG.1

PCT/US99/23253

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			Z/13		
1	gctcacagac (cgggactccg	cctccggttc		ggcgaggcgc
51		ccaacaggtg	cgtgttgtgt		cgcgctccgg
101	gtggagtcaa		gccggcagcc		gggcgggacg
151	gtgccccggg	gcagggctgg	gtggcggccg		gggaggggg
201	ggccgcctcg		ccctggcggc	caatggagac	cgaggccccg
251			cgggggtcag	ccagccttgg	gggccggggc
301	ctggccgggg	acadadaac			cgtccgcgcg
351	gt tataag gc	gagaaattcc	ctgcgccgcg	agccgggagg	cgcacgctcg
401	ctcgtacggc	aaccacaaca	acaaaacaaa		cgggcggcgg
		gcccgggagc	gctCTTCGCT		TCTTGCTCGG
451			TCCCCAGGAC		AGCA T GGGCG
501	ACCTCGGCCA (GCGTCGGGGA (CCCCCCACCC	CCCCACCAC		
551	CTGCCCACCT	CCCGGGACCG	CCACATCCCC	GCGCACGACC	AGCCCGGCGA
601	CIGCCCACCI	TCTGCTGGGA	CCCCCCCCC	CCACATCAGC	CCCTGGGCAC
651	CAAGTGGCTG	GICATCGAGC	CCCCCCTCIA	CCCACCACCG	CGCTGAGGAC
701	AGCGGCACCC	AGGGGGCAGC	CGCCACAACG		
751	GCCACG gtaa	ggaagccata		ccaccggcgg	gtggagcctg
801		gtgggcgtga	tgtcccgctc	cacctgtggg	gccttagcat
851	cctccctccc	ctcgctgacc	tttgacctcc	acgccgggac	ccagagttgg
901	ggtggactag	ccagggccag	atgtggggta	gggagggcag	ttccctgcgt
951	ggaggacccg	cagctgtcca	cggagcaggt	ctgcggggga	ggaggggcc
1001	tcagaggtgg	gtgtgtcatg	ctgcagagcc	tgccctgggt	gaggggctgc
1051		ccaggtccct	gtttcagttc	tgggtcccca	tgctgggtgc
1101		ctaggggtag	ggcagggcag	ggtccccagg	ggccggtaag
1151	gacatgccat	tagaggctgg	gggctgggcc	ggcctgaggt	ctgtggcttt
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1301		agtcacccca		ggcccctggg	gaccccaact
1351	tcccaatccc	agcccctgtc	tagacaggca	gggatgtagc	ctggccccag
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1451		cccacctgca	taataggggt	tggggccacg	
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1601		ccagtcccag	gactgtcggc	gtccctcttg	ccagggccac
1651		ccgattgcca		gttggacaat	cttcactgga
1701		aagaaagccc	ctcttttccc	tttccacccc	atgaagctga
1751	ggagtgagaa		cctgaaattc	taaaaaaaga	aaaaaaaaaa
1801	aaagagaacg	ccttatccat	ggctgttcag	gcgccagacg	ctggcccgag
1851	gggacagcac		tgaagcagcc		atttgagcgt
1901	gcaggtgttt	gcatgtctgg			tgcctttctg
1951	ccagggcgtg			tctccccaaa	ggccttgctg
2001	agreetagee	tcccttcaag	gagtettgtg	gatgcctgct	ctggtctttt
2051	tttaaaaaaa	tatctatttt	atttattatt	atttgtttaa	aaatagagac
2101	aggeteteac	tatottocto	agactagtct	caaagtcctg	ggttcaagca
2151	tteeteetee	ctcagectce	gaaagttctg	ggattacagg	catgagccac
2201	cacteeegge	ctactataat	cttttgtaac	ctagaggaca	grarygarac
2251	agaaaacttt	actcccacc	aaccoccooa	gacagagici	Lycholycoa
2301	agaaaacttt	actoccatoo	caccatctta	gctcactgca	acctccgcct
	cccagactyg	agreeattete	ctacctcaa	ctcccaagta	gctgggatta
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2401	thtasasta	ttaccasacc	tagtatagg	ctcctgacct	cgtgatccac
2451	cccaccatg	ctcccaaact	actagastta	caggcgtgag	ccaccacgcc
2501	ccacctcggc	ngaganagt	tttatttcat	cactotttcc	tgcctggtgc
2551	cggctgggat	acayaaayct	LUCALLUCAL	. Caccagesco	

caggcccatg ctggggttcc tcccaagtgg aattactgac ttaacattta 2601 gcttgggatc ctgagacttc catcacacag ttttctcatt gattcgcagc 2651 caataatatc tgttttaaaa acatctcagg ccgagcgctg tggctcacac 2701 ctgtaatccc agcactttgg gaggctgagg tgggcagatc acctgaggtc 2751 gggagtttga gaccagcctg accaacatgg agaaaccctg tctcttctaa 2801 aaaaatacaa aattagccag gcgtggtggc gcatgcctgt aatcccagca 2851 ctttgggagg ctgaggcagg agaatcgctt gaacccagga gacggaggtt 2901 ccggtgagcc gagatcgcgc cattgcactc cagcctgggc aacaagagca 2951 aaactccgtc tcaaacaaac aaacaaaaaa catctctctg ctccttgggg 3001 ccgggtgcca gctctgctat tggaggcact gagcgacctt gaagcaggca 3051 tgtcactcct ctgtgcccca gtttactcat ctgtaaagtg ggagagctgg 3101 ggcagacagt gagctggctg agggcaggac tgtgtctcct caagcccatg 3151 gcccagggct gccaggtagt agtttgtatt cggtaaatgc tgctggcccc 3201 taagtgtgag cgtgccctgc aaactgcagc gtatggtggg acagccctgc 3251 acggctaccc ctttcctggg tgaccttatt tggttacggt cctatctgaa 3301 gtaggaaagg gacactttag gctgtctctt agctccctca aggccccaca 3351 gcctggacta gagttgccag aaatacttgg tccattcagg ccaaagggac 3401 tgtgaggttg ctgggatggt gcaatcagtc tttgtccatg atgaacccac 3451 agggtagacc aggggttggg ccagcccagt gccctgtgta gttgagccca 3501 ggccccaggc atcccatccc gggcggtggc ctcaggtgga ggtggggcag 3551 ccagttgcca gggatgtgtt ccagcggtca cctctcacca gccccggctg 3601 cccatcagct gttctcaagt ccaggcaatg aagccttcct gccaggaaat 3651 tcccagagtt tctgtgccat gaagtcagcc tgtggccatc ttgggacaca 3701 aggccgggtg ccctggggag agtactctgg gcccttggcc aggtttgtct 3751 gagagtcata ggcagcctga tactagtgga gccagccagg gagggatgag 3801 gcccagccgc tgctggccat aagtatataa gggccatgtg ctgagtgcct 3851 actatgtgcc aggttttgaa atcagtactt gatttattga aaccctctct 3901 tttaatcctc aaggtgcccc tatgaggcac gtaccattta ttgttattgc 3951 cacttgacag atgagaaaac agaggctcag agaggcaaag tggcttgaaa 4001 ttcagtgatt ggtctgggat ttgaatccac agccatgttc ttaagggcat 4051 gctatgctgc cacctatcct gtttatttcc ggcactcatt gattcttcaa 4101 tgtttgactc attaaatcca tcagtgagca tcttctctgt gtcatgcatg 4151 gttctcacct ctgaagatgt agctgtgagc aaaacttcta cagggaatga 4201 gttcacagca gagggatcag ctagagcaaa ggctcagagg tgggaccgtg 4251 cgtcctgtgt tccaggaata cagtatggct gcagcagaga gcagtggaga 4301 gagggcctgg cagtgaggtc tagaggcggc cgggctggct catgctggat 4351 gtttgtgtcc tcggaaggac tttggcttta ttttaaagag gatggggagc 4401 cccagagagc acagcaggga agcctgggga gtctgatgga catttaaaag 4451 gatccttaat ggagagagtg aaggcagagc cttccagaag ggtaagagaa 4501 gggaggatgg agacctgccc tcccccaagg gaggccactc agaagaggta 4551 gagtgtggcc agggcagaga gcaagagagg ctgtggacac aggcacactg 4601 gtccagtgag agccattaga cacattagat ttagcttcat gttgtcttta 4651 gagagggagc cagcctggcc tcgctctatg atcttggaca catcctttca 4701 cttctgggtc tcagtttccc cattagtgtg atgaggatga gaatgctttt 4751 gtcctgggca cactatgagg gtggtgctgg gcacctgggt gcctggttac 4801 catgggcaac aaagctctat tcatgggtgt ggtgaatgca ttgcccacag 4851 caactcaggg cggatgagga gtttcccagc agcccctggt gccctttcgg 4901 ctgaagccct aacaactgtg ggaaaatcca agttccagca gaccccctga 4951 5001 agctggagga gggagtggcc agtgctgcag cagaggctgc ttcatagtaa 5051 ttgcagccaa cagttattga ctaggcactg ttctgagggg tttagatgtg 5101 gtaactgatt gaattcgcct aacaacttta tgaggtaagt cctattgtta 5151 gcccattttg tagatgagga gactgagttt gaaactgggg ggtgtaatgg 5201 aaccttctca ggacccttga agggtagggc ctttgtactc gggccacgag 5251

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5701	gggtatagct	gctttggggc	tactgtgggg	tcagggacac	ttgtgaggcc
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5851	tcgacttggg	ctccgctaca	caggccaaat	ttggatgtcc	catgtttaga
5901	gctgtgtttc	tttgggacct	cttggggcct	cagtttcctc	atctgtaaaa
5951	tgggatactg	atagtgcttc	cccactggcc	tcctctgacg	ggcgccaggg
6001	agaggatggg	acggagcatg	gtgtgctggg	cacgctcctg	ctgtacccac
6051	ccacctggga	gaggggagag	gcaggaatgt	cctgggggtg	tcctttgagg
6101	catagccctg	tcaccccaac	atcctacaaa	ggcatgagaa	ggcagcgagg
6151	acagaccccg	accacctgag	ccctcagcag	ccctgccaca	ctccctgctt
6201	caccccttc	ctgactgatc	tggcacattc	ttgattctcc	tagggagtga
6251	cccaaaatcc	ctccctgccc	tgctgtgtct	ctggggtgga	aggaggctgc
6301	cagcccctcc	tctctcccag	cctcaggctt	ggccaggact	taacaggcag
6351	gcagagaagc	agcttctcca	ctctcttccc	tgacacctgt	aggcccctcc
6401	tgcaggcact	tacctctaag	tggactctca	ggaggaggct	catcagggct
6451	gcagggctca	gaaagagctg	ggctgtggag	ctcttgccaa	ccgccaggcc
6501	ccttctaagt	gctttagcgc	caccgactgc	atcctcccag	cagccttgtg
6551	agatggggat	ttgtggttcc	cagtttactg	atgagaaata	ctgatgagag
6601	atgggtgtgg	tcttgtctgg	ggctccctgg	ctcctggata	gcagctcagg
6651	ttccatcctg	ggcaggctgg	ctctgggaca	ccccccgac	cagctgctgt
6701	gtgggattca	cggtggggct	tgggcagggc	gtgggatctt	ggggccaact
6751	gagccactct	aggcttccag	ggaccaaggc	caggctgagc	tgtctctgta
6801	tcctgagaga	gcatgaacat	cacagaagat	gggcccgggt	tcgaatccca
6851	gctctgccac		gacctgggca	ggggtccctt	cccgctgagc tgggggctgt
6901	cttcatttcc		aatggttcgt	gcccctgctt	agcagctgct
6951	ggagggttgg		cttgttcata	cctgctgttg	tcgctacctc
7001	ctgtgccggc	ctctgaggat	gccactgtga	acagageetg	tttcacccag
7051	caggagcttg	tgtttagggg	tgccgttttg		tccactcgct
7101	ctctgctccg		agagacgtcg		gtagccctgg
7151	tgggtgcgtg				acctcccaca
7201	gtggatgttc				cctgccccc
7251	gttccctgag				tagtgactgg
7301	agcccaggct	cccagcagct			ctgcttcccc
7351	cctcacggca	aggacccccg	cacaccacct		
7401	tctgttccag	gagtggcgac	aagcacagtt		
7451	tcttcacttt	. aagttccggg	aaacgtgcag	aatgtgcagg	
7501	aggtatacat	. gtgccatggt	ggtttgctgc	acccgtcaac gtcctaatgc	
7551	ggttttaago	: tccatataca	ttaggcattt		
7601	cttgcccctc	acccgcccag	taageceegg		
7651	gtgtccatgt	gttctcattg	ttcaactctt		
7701	ctggactctg	atctaacctc	ggtcaaatgg		
7751	gtagcttaac	ctctctgagt	cttagettet		
7801	aggagaggc	cacagagga	caggicacat		
7851	aaggctgttt	gcttccaggt	ceeggeeeg		_
7901	cgcactccct	gatagcatga	gaaycacay		
7951	ctgagagcc	c agectgette	; ccayggaact	gtcacagcco	

ttccccagct ggagccctgt caatggcttt ggggttctct gacacagccc 8001 tgagggggct cacacttccc cttatcattg caaggggtag atctggcttg 8051 aaggccctgg ggcaggcttg gttctgtcct cccctgtcag tgcctcgaca 8101 gggctggcct gggtgaatca ggaccaacgg gaaaggaggc gaggagacca 8151 atctggaccc aagatcctca gctcaataag gtggccccag aactgacatg 8201 gggtgataga gggaagggct gggagggagg agattctggg gccgcagcca 8251 cagettgeac gttgegeegg gtgtgtetgt gegtgeeage tgeatetttg 8301 cgtaccatgt gtgcaaggct gtgtttggct gagtgttcat gtgggccgtg 8351 attgtgggca tgtttctgag tgtctgagtg atgcctgctg gtgtgggctg 8401 gtgggtgtgt ctgcatgtgc gtgtgtgtct ggggagtttc aaaggagaaa 8451 gagggactca ccatcacgct ggctcagcct taaaaaggta ggacatcctg 8501 acacgtgctg caacatggat ggaccttaag gacattgtgc tgagtgaaac 8551 aagccagagg caaaggaaca aacatgtgat ttctcccaga tgaggtttcc 8601 ggaggaggca gatctgtatg gacagaaggt agcatggtgg ttgccggggc 8651 agggggagga gagaatggag aattagtgtt taatggggac agagtttcag 8701 ttggggaagg tgaaaaggtt ctggagctgg atgatggtga tggttggaca 8751 acactgtgca tgcacttaat accactgagc tggacaccta aaaatgctta 8801 caatggtaaa tttcatgtat attttactac aatttttaaa aaattggctg 8851 ggcgtggtgg cttatgcctg taatcccaac actttgggag gccaaggcgg 8901 gaggattgct tgagctcagg agttcaacac cagcctgggc aatatggtga 8951 aaccccgact ctacgaaata tacaaaaatt agcctggtgt ggtggcttgc 9001 acctctaatc ccacctactc agtaggctaa ggcacaagaa tctcttgaac 9051 ctgggaggtg gaggttgcag taagccgaga tcatgccact gcaacccagt 9101 ctgggcgaca gagcaagact ctgtctcaaa aaataaaaga taaataaaaa 9151 aattagaggc caggtgtggc tcacacctgt actctcaaca ctttgggagg 9201 ctgaggtggg aggatcgctt gaagtcaggc atttaagaca tgcctaggca 9251 acatagtgag accttgactc tacaaaaaaa ttcaaaagtt aatgagacat 9301 ggtggcatgt gcctgtagtc ctagctgctg gggaggctga ggtgggagga 9351 tcacttacga ccaggatttc aaggctgcag tgagctgtga ttgcatcact 9401 gcactccagc ctggtgacag agtgaggccc tgtctcaaaa aaatttttca 9451 gtgtttttct gggctgggcg tggtggctca ttcctgtaat tccagcactt 9501 tgggaggctg aggtgggtgg attgcttgag cccaggagtt taagaccagc 9551 tgggcaacat ggcaaacctc atctctacaa aaaataaaaa taaaaaatta 9601 gctgggcatg gtggtgcaca cctgtactaa cagctacgag agaggctaag 9651 gtgggaggat cacctgagcc cgggaggttg aggctgcagt gagccatgat 9701 tgcaccactg cactctagcc tgggcgatac agcaagaccc tatctcaaaa 9751 aaaaaaaaa aaaaaaaaa aaaaacaccc agtggggtca gtagaacccc 9801 aagagtette tteeeteea geteeetgt acaccageee cagetetgea 9851 ggtagctggg ggcccagaca gcttcctggg gacccccagc cttccctctg 9901 ccctttttc taccagtttt gctgcccctc cttcaagact catgtccaga 9951 gggggtgaga tctgcactta tacagccccc tcctctgtaa tgagtgagcc 10001 aagtcagccc aggttattcc agaaggggca ccctaccagc ccccagtcc 10051 ccaagctgcc ctgggcctat aaaagcaggc aaggggaccc ctagtagatc 10101 atgtaggtgt tacctcttag tgggtgctgg aggggcctga agtgctttct 10151 tececeaggg tggtaggaga atgteetgge agtgaettea gggeeegetg 10201 tcacttccgt tttaagactc accagctggt aggctcatta gcaagaggac 10251 aataggaggc ccctgtcctc agtcagcttt cttcaaaggt gtttccttta 10301 gcaactggga ggcctccctt ctccagaccc atggggacaa caccacccag 10351 ctactggttc tataagctgc tgtatggctc tggctagccc attcagagaa 10401 agcetetgaa agtacaagga aaaaaatcag tecaagaget gtgaacaatt 10451 agtgagccga ttacaatacc aagaccacag gcagacctgg aaggctaagt 10501 gageceaggt gtgaagttea agettaettt acttetggge caetteetgg 10551 ctggtctctt tecctggccc ttatetttet cetggtctgt ettetettet 10601 caccccttt ctttactctt tcttccttct cctgcatcgt actccacccc 10651

WO 00/21557

6/19 cactccagct attacacaga atcgcgagaa tgttggatta ttcattttat 10701 ttatgatgtt ttctttttig taaaaataga gacaaggtct cactatgtgg 10751 cccaggctgg tettgaacte etggcetcaa gcaateeteg tgeettggee 10801 tcttacagtg ctgggattac agatgtgagc caccatgcct ggcccatttt 10851 atttacttta aaaaaaaat taggctgggc gcggtggctc acacctataa 10901 ttccagcact ttgggaggcc aaggtgggca gatcaactga ggtcaggagt 10951 taaagaccag cttggccacc tggggtcagg agtttgagac cagctactcc 11001 ggaggctgag accggagaat tgcttgaacc caggaggtag aggttgcaat 11051 gaactgagat catgccattg catgccagcc tgggcaacag agcaagactg 11101 totcaaaaaa aaaaaaaati atgittigig ofootgotto ofgottigta agtcaaatca gittaactgi toaagtgici toottgoaaa ooccoaagga 11151 11201 ctcaatgtgt gtcgcccttg actgatcccc ccgccccgtg acccagtggt cctcagttcc aggttttccc acctacctt cacccactgc ttatgtttat 11251 11301 aaaaacgggg taaatcaaat gttcgtgacc cagatcttat tctacatgca 11351 gtggaaactt gtatgactta agctttttgg aaaagcagaa ccttttttcg 11401 tggttcaaga aatcaaagtc ttcccgggag gtctttctgt aaatccagag 11451 ctgcagatgt ttgaccgtgt tcagagaggg gcccttgtgc tgggtgaagt 11501 ggatggggca cagcaggcaa tgggtgaaaa gcaggacaac ctggggccct 11551 gggaggacca gggagggccc atgtctttga ctgttcatca gccggctgac 11601 ttcctgtccg cctgtcgtct gctctgccca tccatccgta gtccttccgc 11651 ctgtctctgc tggttgccgc tgtgctactc agctgtgtct gtctgtccgc 11701 ctgactgtct gctctccttc aggatgcctt ccgtgccttc catcaagatc TCAATTTTGT GCGCAAGTTC CTACAGCCCC TGTTGATTGG AGAGCTGGCT 11751 11801 CCGGAAGAAC CCAGCCAGGA TGGACCCCTG AATgtgagcc agagccctag 11851 gagaggctca gcccctgagg gagggggatg gctggagggc tgggagacat 11901 tgccacatgg ccaggagcag ctccctcggc attcgcccaa ggggatgcag 11951 agccagggct gagcctgccc tcccctccca gggggcaggc agttgaaagt 12001 gaagctgtag ggatgccctg agaagtccag ggctccagat ctggtttagc 12051 caggcactcg titggatecc gaggcaaget cectecetgt tgtegeecag 12101 tgtccccatc aaaaggagga ttttgatgaa ctgatttctc tcctggctgt 12151 agcgtcttac ccaccccata ccttttggga gggagaggag gcttcaccac 12201 cagccagtgc tccagctcac accccgggct gggtactctt gtcacttcat 12251 tectetttge ceacaceet tgggeetgge gatgggagga geggetgggg 12301 ctccaggaga atgggggtgg ggaggaattt cttccttggc tgatcggccc ctctgctatg gcagGCGCAG CTGGTCGAGG ACTTCCGAGC CCTGCACCAG 12351 12401 GCAGCCGAGG ACATGAAGCT GTTTGATGCC AGTCCCACCT TCTTTGCTTT 12451 CCTACTGGGC CACATCCTGG CCATGGAGGT GCTGGCCTGG CTCCTTATCT 12501 ACCTCCTGGG TCCTGGCTGG GTGCCCAGTG CCCTGGCCGC CTTCATCCTG 12551 GCCATCTCTC AGgtgacccc agttctgtgt tgcagccacc ttaactgccc 12601 aacagacgtg ggcccccatg catctgggca ttgtgaacat atttgctaaa 12651 tgaatgaatg gacctatgaa aggatgaatg gatgaataaa cagatgaatg 12701 agtgaacagt ctgaaggccc atcaggcatg tctgtgggtc aagctgcatt 12751 ccagatgage caagaagtte ettettgaac agatteegat caageacagg 12801 gccactgagc cagaggctgc tgccctgcag cttcatgaca cttacgagcc 12851 cctccacctc cctgggactc agttctcatc tgtaaaaaga ggacactggc 12901 ccacaagggt cttgaaatgg agcattagca cggggggtacc ctgcaagctg 12951 aaaggattca ctggggcccc aggccctggc gggctccgtc cttcccaaca gcttctgacc ctgcctctct ccccagGCTC AGTCCTGGTG TCTGCAGCAT 13001 13051 GACCTGGGCC ATGCCTCCAT CTTCAAGAAG TCCTGGTGGA ACCACGTGGC 13101 CCAGAAGTTC GTGATGGGGC AGCTAAAGgt gagggtgggg tgggtggtca 13151 gccaggtgct gggtggcgct gggtctgccc aagtgtgtgg gcacagtcgg 13201 gggcacagec tgccctgaga gccccctcct cctccacagG GCTTCTCCGC 13251

FIG.2E

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			•			
	13301	CCACTGGTGG	AACTTCCGCC	ACTTCCAGCA	CCACGCCAAG	CCCAACATCT
•	13351	TCCACAAAGA	CCCAGACGTG	ACGGTGGCGC	CCGTCTTCCT	CCTGGGGGAG
•	13401	TCATCCGTCG	AGgtgggtgg	ggagggacct	ggacaacctc	tggctgggcc
•	13451	tgcagctgag	ggggagctaa	tgcactgggt	ccccactctg	cccctgacct
	13501	agcccctgat	ctggcctcca	ctctggctgg	gccaagctct	gcccccgtgt
	13551		cacctcccaa		acgaccagcc	cgcttgctag
	13601	aatctagagt	tgcctttgac	ccttggcccc	agccagcccc	gtgaccttgc
	13651		gaggtggcct	ggagagctgc	tgtctccagc	cgccgcctgt
	13701	ctccacagTA	TGGCAAGAAG	AAACGCAGAT	ACCTACCCTA	CAACCAGCAG
	13751	CACCTGTACT	TCTTCCTGAg	tgagtgtcca	tctgtccttc	tggggtgggg
•	13801	gagtgcctgg	gcctgcactg	tcctccctgc	tgtcctggac	cactcccagc
	13851	cacttcctgg	ggcggggcac	gtctgtcagg	tctccctggt	catggcatcc
	13901		tgcagtctgt	acacactctc	ccagcagcat	gcctttgccc
	13951	cagctgtctc	ccgtgcctgg	gacaccttgc	agccacgggc	catcacagcc
	14001	ctgctgggag	cttccccaag	ccccacgtag	aatttcttct	tgccctcact
	14051	agagtggtcc	ggagccctag	agtctttggg	cagttgttgg	ggcggacaga
	14101	gtgaggactc	aagtctggcc	ctgacttgcg	gtgaagggtg	gtgggaggtg
	14151	gtggggtaag	ggcagcctgg	ggaggcttgg	acacagaatt	gggggtgata
	14201	tggggtcatt	cagctggatg		caacgtccca	ggggcattcc
	14251	tggagtaaca	gagcccctca		cactcacctt	ggcagcccag
	14301	ccccactcct	gaacactctc	atgccccttc	ttgcag <u>TCGG</u>	CCCGCCGCTG
	14351	CTCACCCTGG	TGAACTTTGA	AGTGGAAAAT	CTGGCGTACA	TGCTGGTGTG
	14401	CATGCAGTGG	GCG gtgagtg	gggttgccca	ggaccccggg	catacggctg
•	14451	ccgtggcagg	aggtggtgcc	tcgggggaca	gtacctgccc	atgaaggcaa
	14501	acagggtgca	catgtgcgtg	caacagtgtg	gctcacatgt	atgcgtgcaa
	14551	cagtgtggct	cacatgtgtg		gagagcgagt	gtgcccgtga
	14601	ctgtacgtgt	ggtgggggg		cagggggggt	gtgggtctct
	14651	ctcggtgagg	gtgtcttccc	aggaggagtt	gctgggccga	ctctgccagg
	14701	catctgtgtc	cctggcaggg		cacaccctgc	atgacacctt
	14751	cgtcactaaa	atcagcctcg	tgagctggca	gggcaaggac	cctgttcctt gggctctgag
	14801	tactcagctg			ggcctgtcct	
	14851	gcaaatcagg	cagaagggtt	ggatgcctga		gcctctgccc
	14901	ggcctccaga	cctccgggca	cctggagacc	tctcggtatc CTTCTATGCC	
	14951	tcctctgcag	GATTTGCTCT	GGGCCGCCAG		
	15001	TATCCTACCT			GGGTGCTGCT	
	15051	<u>GCTGTCAG</u> gt	atggcaggga	gtggcgaggt	cacacacagg	
	15101	ccccactgc		cagagcttcc	cttttcccgt	ctgcagaatg
	15151	gggccagtgg	tactgcctcc	ctggcttgct	ggtggaatca	
	15201	agcgtggcag		tcggtgggtt	tagggagcgt	
	15251	gtaagtggcc		cggagctgct		
	15301	gacactgctc	cattcagatt	ctttaaacac	tggcaagggg	
	15351	caatcctatt	gtacagataa	ggaagtcaag	gccactuggg	gacagctgct
	15401	ctccagcctc	cactcagggt	gcctaagtgg	CCACTGGTTC	ctagggcagt crcrccarca
	15451	gcccgagcct	ccccacag <u>GG</u>	TCCTGGAAAG	CCCACCACAA	GTGTGGATCA
	15501	CACAGATGAA	CCACATCCCC	AAGGAGATCG		GCACCGGGAC
	15551	TGGGTCAGCT	CTCAGgtggg	cagcaggggt	ggggccatc	ctgggtgggg
	15601	tggggggtcc	cagctaggag	ccagatggca	aagcagggat	gaggccctga
	15651	agaggetace	aggragaga	taataccata	r adatcaqqqa	LCLGCaacgg
	15701	cotoctcaca	tataccccac	caacttccaa	cauciggene	CCUC CONTRACTOR
	15751	CGTGGAGCCC	TCACTTTTCA	CCAACIGGIII	CAGCGGGGAC	CICAACAACA
	15801	AGATCGAGCA	<u>CCA</u> gtgagtg	rgggtgctgg	gggccagugg	gaggtgggga
	15851	gggggtcctg	ggagggato	ctgggagggg	acccgtgggt	ggggcctctc

FIG.2F

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				8/19		
	15901	tctggaatct	cccacttcag	gtgccagcat	acgctcccca	ccccag <u>CCT</u>
	15951	CTTCCCCAGG	ATGCCGAGAC	ACAACTACAG	CCGGGTGGCC	CCGCTGGTCA
	16001	AGTCGCTGTG	TGCCAAGCAC	GGCCTCAGCT	ACGAAGIGAA_	GCCCTTCLAC
_	16051		TGGACATCGT	CAGgtgaggc	tgcagcccgg	ccctctgtt
_	16101	ctggtggctt		tatgcctacc	cttgtccagg	tcagcctcat
	16151	gctgagcccc		gagcctttct	gtccacgtcc	catgcccttc
	16201	ctcccttccc	cagccttcac	gcacacagtg	agaatttctg	gagcacctac
	16251	tgcagactca		tgcctgcggt	gagcaggtct	atgcaaacct
	16301	accccaaag	actaagggaa	aaaagctaac	agatccagtt	tctcagaagg
	16351	accectaa	cagggactca	taaacagaag	ccatgtctca	gggccgggtg
	16401	cggtggctca	cgcctgtaat	tccagcactt	ggggaggctg	aggtgggcgg
	16451	atcacttgag	gtcaggagtt	cgagaccagc	ctggccaaca	tggtgaaacc
	16501	ccgtctctac	taaaaaaaaa	aaaaaaaaac	aaaacaaaac	aaaaattagc
	16551	tgggtgtggt	ggcaggtgcc	cataatccca	gctacttggg	aggctgaggg
	16601	aggagaatca	cttgaactcg	caggggcaga	ggttgcagtg	agctgagatt
	16651	gtgcctttgc	agtccagcct	gggcaacaga	gcaagactct	ctcaaaaaca
	16701	22222222	ccatgtctca	ggcagccaag	agttgggaca	tcccctcaca
	16751	cgccctctag		ctatatagca	agcttttagg	gtgaacccca
	16801	tgcaggtggt	tcttatgaac	ctggtgacca	ctggaggtta	gataagcgtc
	16851	tacaagagga	ggttatctat		tggcattcag	ggtcaagcat
	16901	cggtcatcag		ttgaagatgg	cattgccctt	gtagcaatgc
	16951		gagcttcctg	ccctcttgga	gctgatgttc	cttccagcaa
		aggetetaga	aagcaattaa	aataacaaat	aagtacatta	cagaagatgg
	17001 17051	aggaaacagc gcaaaagaac	aatgaaaagc	ccctcagggt	ggggacaggg	gaggggaggg
	17101	_	gcaggggcgg	cagtttctaa		gggtgggcag
		gggcggccag	ctgacgtgtg	agcagggaca	gggaggaggg	gagaggtctc
	17151	tattgacagg	catctggcaa	agagcgttca	ggcagagggc	acttgaccct
	17201	gccacaggga	ctcatggcat	agatagccga	ggcaggcatg	caggcactca
	17251	gaatgccaag	acgcccggct	tgcatcttgg	aaagctgccc	ctactgggaa
	17301	gagaagggac	gcaggagtcg	aagtggaaaa	ggagagcaga	ggacactgca
	17351	tgactggcgg	caaggagteg	tggggctcag		caccttggag
	17401	gccatccagg		tccaggtctt	atacctctgc	gcctttgtac
	17451	gtggggaaca	ccttacttgg		ttcctgtgct	ggtgttcaga
	17501	acgctgttcc	tectteatga	tctctcccag	cctgatgctc	
	17551	tgcccacttc				cctagcagat
	17601 17651	ccatttggca tgttgacatt	teteetee	ctgcccaata		gatcgggtgg
	17701	gcaggttcca			tgcagcgccc	agcaggaggc
	17751	agcaatggag	_			ggctcatgcc
	17801	tggacttggc		ctccagctcc		acccgtcacc
	17851	ccggtctaga		gagaatgagc		tctcccaacc
		as acat acaa	coccetecc	tacctacccc	cagggaaggg	aacccacagg
	17901 17951	gastggggst	ctccactcac	acttaccato	ggggatacag	gggtgttagg
					CCACLUCCAC	CCCCACCCC
	18001	accurrence and accurrence and accurrence accurrence and accurrence	AAGAAGTCTG	GTGACATCTG	GCTGGACGCC	TACCTCCATC
	18051	20001CCCIG	ACACCCACC	CCCAGAGA	GGGCTCAGG	CTCCCCAGCAAC
_	18101	AGT GAAGGC	CCCCCCCCC	TUCATACCO	CACCCCTCCA	CTGGCCAGCC
_	18151	CAAGCCAGCC			CHACACAAC	CCTCGGCCCC
_	18201	TGGGGGTGCC	CIGCCIGCCC	TCCTGGTWCT		GCCTGATGGG
_	18251	CTCACATGTG	TATTCAGCAG	CCCTATGGCC		GCCTGATGGG ACCGAGAATT
_	18301	ACAGGGGTAG	AGGGAAGGTG	AGCATAGCAC	M ACAMMONACA	AGCGAGAATT
_	18351	GGGGGAAAGC	TGTTATTTT	' ATATTAAAA	T ACATICAGA	T GTATTATGGA
_	18401	GT				

FIG.2G

1	CTTCGCTTCCCTCGGGGTCTTGCTCGGACCTCGGCCACCGCCTGGGATCC	50
51	CCAGGACTCGTGCGTGCAGCATGGGCGGCGTCGGGGAGCCGGGACCGCGG	100
1	M G G V G E P G P R	10
101	GAGGGACCCGCGCGGGGGCACCGCTGCCCACCTTCTGCTGGGAGCA	150
11	E G P A Q P G A P L P T F C W E Q	27
151	GATCCGCGCGCACCAGCCCGGCGACAAGTGGCTGGTCATCGAGCGCC	200
28	I R A H D Q P G D K W L V I E R R	44
201	GCGTCTACGACATCAGCCGCTGGGCACAGCGGCACCCAGGGGGCAGCCGC	250
45	V Y D I S R W A Q R H P G G S R	60
251	CTCATCGGCCACCACGGCGCTGAGGACGCCACGGATGCCTTCCGTGCCTT	300
61	L I G H H G A E D A T D A F R A F	77
301	CCATCAAGATCTCAATTTTGTGCGCAAGTTCCTACAGCCCCTGTTGATTG	350
78	H Q D L N F V R K F L Q P L L I G	94
351	GAGAGCTGGCTCCGGAAGAACCCAGCCAGGATGGACCCCTGAATGCGCAG	400
95	E L A P E E P S Q D G P L N A Q	110
401	CTGGTCGAGGACTTCCGAGCCCTGCACCAGGCAGCCGAGGACATGAAGCT	450
111	L V E D F R A L H Q A A E D M K L	127
451	GTTTGATGCCAGTCCCACCTTCTTTGCTTTCCTACTGGGCCACATCCTGG	500
128	F D A S P T F F A F L L G H I L A	144
501 145	CCATGGAGGTGCTGGCTGGCTCCTTATCTACCTCCTGGGTCCTGGCTGG	550 160
551 161	GTGCCCAGTGCCCTGGCCGCCTTCATCCTGGCCATCTCTCAGGCTCAGTCVPSALAAFILAISQAQS	600 177
601 178	CTGGTGTCTGCAGCATGACCTGGGCCATGCCTCCATCTTCAAGAAGTCCTWCLQHDLGHASIFKKSW	650 194
651 195	GGTGGAACCACGTGGCCCAGAAGTTCGTGATGGGGCAGCTAAAGGGCTTC W N H V A Q K F V M G Q L K G F	700 210

701	TCCGCCCACTGGTGGAACTTCCGCCACTTCCAGCACCCACGCCAAGCCCAA	750
211	S A H W W N F R H F Q H H A K P N	227
751	CATCTTCCACAAAGACCCAGACGTGACGGTGGCGCCCGTCTTCCTCCTGG	800
228	I F H K D P D V T V A P V F L L G	244
801 245	GGGAGTCATCCGTCGAGTATGGCAAGAAGAAACGCAGATACCTACC	850 260
851	AACCAGCAGCACCTGTACTTCCTGATCGGCCCGCCGCTGCTCACCCT	900
261	N Q Q H L Y F F L I G P P L L T L	277
901 278	GGTGAACTTTGAAGTGGAAAATCTGGCGTACATGCTGGTGTGCATGCA	950 294
951	GGGCGGATTTGCTCTGGGCCGCCAGCTTCTATGCCCGCTTCTTCTTATCC	1000
295	A D L L W A A S F Y A R F F L S	310
1001	TACCTCCCTTCTACGGCGTCCCTGGGGTGCTGCTCTTCTTTGTTGCTGT	1050
311	Y L P F Y G V P G V L L F F V A V	327
1051	CAGGGTCCTGGAAAGCCACTGGTTCGTGTGGATCACACAGATGAACCACA	1100
328	R V L E S H W F V W I T Q M N H I	344
1101 345	TCCCCAAGGAGATCGGCCACGAGAAGCACCGGGACTGGGTCAGCTCTCAG P K E I G H E K H R D W V S S Q	1150 360
1151	CTGGCAGCCACCTGCAACGTGGAGCCCTCACTTTTCACCAACTGGTTCAG	1200
361	L A A T C N V E P S L F T N W F S	377
1201	CGGGCACCTCAACTTCCAGATCGAGCACCACCTCTTCCCCAGGATGCCGA	1250
378	G H L N F Q I E H H L F P R M P R	394
1251	GACACAACTACAGCCGGGTGGCCCCGCTGGTCAAGTCGCTGTGTGCCAAG	1300
395	H N Y S R V A P L V K S L C A K	410
1301	CACGGCCTCAGCTACGAAGTGAAGCCCTTCCTCACCGCGCTGGTGGACAT	1350
411	H G L S Y E V K P F L T A L V D I	427
1351	CGTCAGGTCCCTGAAGAAGTCTGGTGACATCTGGCTGGACGCCTACCTCC	1400
428	V R S L K K S G D I W L D A Y L H	444

WO 00/21557 PCT/US99/23253

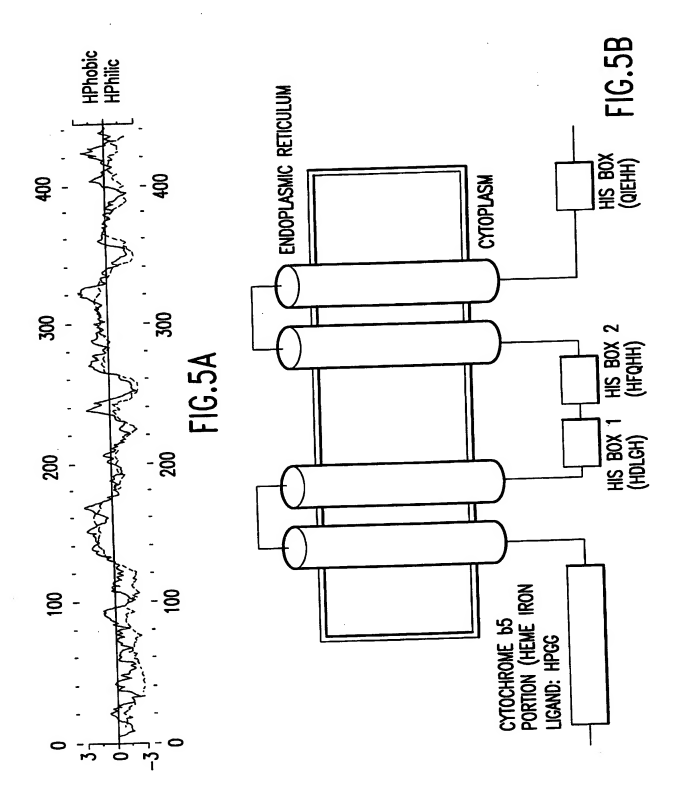
11/19

L401 445	ATCAGTGAAGGCAACACCCAGGCGGGCAGAGAAGGGCTCAGGGCACCAGC Q	1450 445
1451	AACCAAGCCAGCCCCGGCGGATCGATACCCCCACCCCTCCACTGGCCA	1500
1501	GCCTGGGGGTGCACTGCCTGCCCTCCTGGTACTGTTGTCTTCCCCTCGGC	1550
1551	CCCCTCACATGTGTATTCAGCAGCCCTATGGCCTTGGCTCTGGGCCTGAT	1600
1601	GGGACAGGGTAGAGGGAAGGTGAGCATAGCACATTTTCCTAGAGCGAGA	1650
1651	ATTGGGGGAAAGCTGTTATTTTATATTAAAATACATTCAGATGTAAAAA	1700

FIG.3C

1	GTACAGCGGCAATGGGCGGTGTCGGGGGGGACCCGGAGGGGGGACTCGGGCCG M G G V G E P G G G L G P	50 13
51	CGGGAGGGCCCGCACCGCTGGGGCGCCCCTACCCATCTTCCGCTGGGA	100
14	R E G P A P L G A P L P I F R W E	30
101	GCAGATCCGCCAGCATGACCTACCAGGCGACAAGTGGCTGGTCATCGAGC	150
31	Q I R Q H D L P G D K W L V I E R	47
151	GCCGTGTCTACGACATCAGCCGCTGGGCACAGCGGCACCCAGGGGGGTAGC	200
48	R V Y D I S R W A Q R H P G G S	63
201 64	CGCATCATCGGCCACCACGG 220 R I I G H H 69	

FIG.4



PROFILESCAN of : CYB5rp_correct_protein check: 5714 from: 1 to: 445

```
GETSEQ from bmd, December 2, 1997 14:20.
Compare to profile library: GenRunData:profilescan.fil
Profile: profiledir:cytochrome_b5.prf
                         Gap Length weight: 0.05
   Gap weight: 4.50
                         Ave mismatch
                0.27
   Ave match:
(Peptide) PROFILEMAKE v4.40 of: 0191.Msf2{*} Length: 48
  Sequences: 24 MaxScore: 27.58 December 2, 1992 00:07
This profile is derived from PROSITE release 10.0 and has been tested
by a database search against SWISS-PROT release 26.0. A comparison
of the SWISS-PROT annotation and the results of the database search follows.
For further information about this motif, consult the . . .
Profile: profiledir:cytochrome_b5.prf alignment: 1
                   Gaps: 0
 Quality: 20.77
    Ratio: 0.43 Length: 48
 Normalized quality: 2.91
      31 HDQPGDKWLVIERRVYDISRWAQRHPGGSRLIGHHGAEDATDAFRAFH 78
S
          |: ..: ||||. .|||:::| . ||||. | . .||.|:||.| ::|
       1 HNDGEETWLVVNGQVYDITKFLEEHPGGPDVIMEAAGTDATEEFEAIH 48
*Cytochrome b5 family, heme-binding domain signature *
```

FIG.6

```
① pir:s68358 hypothetical protein - common sunflower
 Length = 458
Score = 169 (79.4 bits), Expect = 2.8e-42, Sum P(4) = 2.8e-42
 Identities = 31/85 (36%), Positives = 49/85 (57%)
                                                     His box 3
         348 IGHEKHRDWVSSQLAATCNVEPSLFTNWFSGHLNFQIEHHLFPRMPRHNYSRVAPLVKSL 407
Query:
                             T ++ S + +WF G L FQ+EHHLFPR+PR +
                                                                 ++P+ + L
             +G K +W
                        Q
         348 VGPPKGDNWFEKQTRGTIDIACSSWMDWFFGGLQFQLEHHLFPRLPRCHLRSISPICREL 407
Sbjct:
         408 CAKHGLSYEVKPFLTALVDIVRSLK 432
Query:
                         F A V +++L+
             CK+LY
         408 CKKYNLPYVSLSFYDANVTTLKTLR 432
Sbjct:
 Score = 133 (62.5 bits), Expect = 2.8e-42, Sum P(4) = 2.8e-42
 Identities = 21/53 (39%), Positives = 35/53 (66%)
                                              HPGG motif
          26 EQIRAHDQPGDKWLVIERRVYDISRWAQRHPGGSRLIGHHGAEDATDAFRAFH 78
Query:
             ++++ H+ P D W+ I +VY+++ WA+ HPGG
                                                       +D TDAF AFH
                                                 + +
          22 KELKKHNNPNDLWISILGKVYNVTEWAKEHPGGDAPLINLAGQDVTDAFIAFH 74
Sbjct:
 Score = 118 (55.5 bits), Expect = 2.8e-42, Sum P(4) = 2.8e-42
 Identities = 25/76 (32%), Positives = 34/76 (44%)
                                                                  His box 2
                                   His box 1
         165 LAAFILAISQAQSWCLQHDLGHASIFKKSWWNHVAQKFVMGQLKGFSAHWWNFRHFQHEA 224
Query:
                                                       + G S WW + H H H
                           L HD GH +
                                           WN A F+
             L+ IL ++ Q
         152 LSGAILGLAWMQIAYLGHDAGHYQMMATRGWNKFAGIFIGNCITGISIAWWKWTHNAHHI 211
Sbjct:
         225 KPNIFHKDPDVTVAPV 240
Query:
               N
                    DPD+
         212 ACNSLDYDPDLQHLPM 227
Sb jct:
 Score = 34 (16.0 bits), Expect = 2.8e-42, Sum P(4) = 2.8e-42
  Identities = 7/14 (50%), Positives = 9/14 (64%)
```

FIG. 7A

```
gp:bou79010 1 PID:g2062403 Borago officinalis delta 6 desaturase mRNA,
 complete cds. (gb:U79010) (NID:2062402)
 Length = 448
Score = 179 (84.1 bits), Expect = 2.3e-42, Sum P(3) = 2.3e-42
Identities = 34/87 (39%), Positives = 48/87 (55%)
                                                    His box 3
        348 IGHEKHRDWYSSQLAATCNVEPSLFTNWFSGHLNFQIEHHLFPRMPRHNYSRVAPLVKSL 407
Query:
                                    + +WF G L FQIEHHLFP+MPR N +++P V L
                        Q
                            T ++
             +G K +W
        338 VGKPKGNNWFEKQTDGTLDISCPPWMDWFHGGLQFQIEHHLFPKMPRCNLRKISPYVIEL 397
Sbjct:
        408 CAKHGLSYEVKPFLTALVDIVRSLKKS 434
Query:
                        F A
                                 +R+L++
             CKHLY
         398 CKKHNLPYNYASFSKANEMTLRTLRNT 424
Sbjct:
 Score = 144 (67.7 bits), Expect = 2.3e-42, Sum P(3) = 2.3e-42
 Identities = 23/53 (43%), Positives = 36/53 (67%)
                                              HPGG MOTIF
          26 EQIRAHDQPGDKWLVIERRVYDISRWAQRHPGGSRLIGHHGAEDATDAFRAFH 78
Query:
             ++++ HD+PGD W+ I+ + YD+S W + HPGGS +
                                                      ++ TDAF AFH
          12 DELKNHDKPGDLWISIQGKAYDVSDWVKDHPGGSFPLKSLAGQEVTDAFVAFH 64
Sbjct:
 Score = 105 (49.3 bits), Expect = 2.3e-42, Sum P(3) = 2.3e-42
 Identities = 22/68 (32%), Positives = 28/68 (41%)
                                                              His box 2
                         His box 1
         176 QSWCLQHDLGHASIFKKSWWNHVAQKFVMGQLKGFSAHWWNFRHFQHHAKPNIFHKDPDV 235
Query:
                                           LGS WW + H HH
                                       F
             QS + HD GH +
                              SN
         153 QSGWIGHDAGHYMVVSDSRLNKFMGIFAANCLSGISIGWWKWNHNAHHIACNSLEYDPDL 212
Sbjct:
         236 TVAPVFLL 243
Query:
                 p ++
         213 OVIPFLVV 220
Sbject:
```

FIG. 7B

```
①pir:s35157 Delta(6)-desaturase - Synechocystis sp.
Length = 359
```

Score =126 (59.2 bits), Expect = 9.0e-09, Sum P(2) = 9.0e-09Identities = 21/54 (38%), Positives = 33/54 (61%) His box 3

372 FTNWFSGHLNFQIEHHLFPRMPRHNYSRVAPLVKSLCAKHGLSYEVKPFLTALV 425 Query: F NMF G LN Q+ HHLFP + +Y ++ ++K +C + G+ Y+V P

292 FWNWFCGGLNHQVTHHLFPNICHIHYPQLENIIKDVCQEFGVEYKVYPTFKAAI 345

Sbjct:

Score = 36 (16.9 bits), Expect = 9.0e-09, Sum P(2) = 9.0e-09Identities = 6/15 (40%), Positives = 8/15 (53%) His box 2

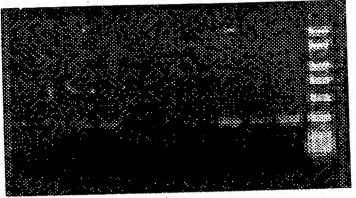
209 GFSAHWWNFRHFQHH 223 Query:

G S+ W +RH

113 GLSSFLWRYRHNYLH 127 Sbjct:

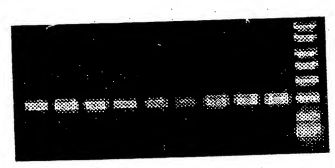
FIG.8





- 2.
- Heart Brain Placenta
- Lung
- Liver
- Skeletal Muscle Kidney Pancreas 6. 7.
- 8.
- Retina

FIG.9A



LPCR Marker

- Heart
- **Brain** 2.
- 3. **Placenta**
- Lung Liver

- Skeletal Muscle Kidney Pancreas Retina 6.
- 7.
- 8.
- 9.

FIG.9B

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INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/23253

A. CLASSIFICATION OF SUI	SJECT MATTER		1			
IPC(7) :A61K ^{29/395} ; C12P 7/62; C12N 9/02, 15/00; C07H 19/00 US CL :435/135, 189, 320.1, 452.3; 424/130.1; 536/23.2						
According to International Patent Cl	According to International Patent Classification (IPC) or to both national classification and IPC					
B. FIELDS SEARCHED						
Minimum documentation searched (y classification symbols)				
U.S. : 435/135, 189, 320.1, 452	2.3; 424/130.1; 536/23.2					
Decrease seembed other than n	ninimum documentation to the ex	stent that such documents are included	in the fields searched			
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Please See Extra Sheet.						
Electronic data base consulted durin	g the international search (name	of data base and, where practicable,	search terms used)			
Medline						
Search terms: CYB5RP, delta-6, fa	tty acid desaturase, human or n	omo sapiens.				
C. DOCUMENTS CONSIDER	ED TO BE RELEVANT					
Category* Citation of docume	nt, with indication, where appro	priate, of the relevant passages	Relevant to claim No.			
X Database GenE	Bank, Accession AA	C23396, submitted by	1-15			
LAMERDIN, JE	, publicly available on	12 June 1998, see entire				
record.		·				
1		2004770 submitted by	1-15			
X Database Genl	Bank, Accession AC	C004770, submitted by 12 June 1998, see entire	1-13			
LAMERDIN, JE	y identification of CDS	at about line 50.				
record, especiali	y identification of CDD					
X,P Database GenBa	nk, Accession AAD312	282 submitted by LI et al,	1-15			
publicly available	e on 19 May 1999, see	e entire record.				
	A CHINANI CENON	AE SCIENCES INC) 11	1-15			
X WO 98/39446	A2 (HUMAN GENUN	ME SCIENCES, INC.) 11 especially SEQ ID No:63.	• • •			
September 1998.	, see entire document, c	specially 52Q 12 1.6.65				
			}			
		·				
Further documents are listed	in the continuation of Box C.	See patent family annex.				
Special categories of cited docur	nene.	To later document published after the in date and not in conflict with the ap	DISCRIPTION OUT CITED IN CITEDERIC			
"A" document defining the general st to be of particular relevance	ate of the art which is not considered	the principle or theory underlying u	ie maeurion			
B carlier document published on o	r after the international filing date	document of particular relevance; to considered novel or cannot be considered.	ne claimed invention camer be lered to involve an inventive step			
and the same and which may throw do	ubts on priority claim(s) or which is	when the document is taken alone 'Y' document of particular relevance; t	he claimed invention cannot be			
special reason (as specified)		considered to involve an inventor	ch documents, such combination			
means ·	disclosure, use, exhibition or other	being obvious to a person skilled in	tie art			
the priority date claimed						
Date of the actual completion of the	ie international search	Date of mailing of the international so				
24 FEBRUARY 2000		15MAR/2000				
Name and mailing address of the		Authorized officer	& plan. Jon			
Commissioner of Patents and Trad Box PCT	INIT	BRADLEY S. MANTEW	$\overline{}$			
Washington, D.C. 20231		Telephone No. (703) 308-0196	$\mathcal{O}_{\mathbb{R}}$			

INTERNATIONAL SEARCH REPORT

International application No. PCT/US99/23253

B. FIELDS SEARCHED Documentation other than mini	mum documentation that are included in the fields s	earched:
Because a CRF was not made appears to encode the same de- available amino acid and nucle	available at the time of the search, Database GenBa saturase as set forth in Figures 3A-C of the instant a tic acid databases.	nk Accession AF134404, which application, was scarched against all